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(54) **Method of cleaning substrates**

(57) A method of cleaning electronic components, characterized by cleaning with an oxidizing cleaning fluid, followed by cleaning with a reducing cleaning fluid with the application of ultrasonic vibrations.

This provides a simple cleaning method which can

remove metal, organic and fine particle contaminants on the surface of electronic components, and especially those on silicon bases, and also suppress an increase in the roughness of base surface on the order of atoms during cleaning processes.

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Description

[0001] This invention concerns a method of cleaning electronic components. More specifically, this invention pertains to a method of cleaning electronic components that is capable of effectively removing contaminants, such as fine particles which adhere to the surface, without making the surface of the base rough in the wet cleaning process used in industries in which such electronic components as semiconductor silicon bases are handled.

[0002] Components such as silicon bases for semiconductors and glass bases for liquid crystals have conventionally been cleaned by RCA cleaning, a heat cleaning method that uses hydrogen peroxide-based concentrated chemical solutions (sulfuric acid + hydrogen peroxide water, hydrochloric acid + hydrogen peroxide water + water, ammonia + hydrogen peroxide water + water). However, wet cleaning processes began to receive attention in recent years for reasons pertaining to environmental protection and resource conservation.

[0003] Against this backdrop, the inventor, et al. earlier proposed a method of cleaning with an oxidizing cleaning fluid, such as ozone water, to remove mainly metal and organic contaminants, and another cleaning method using a reducing cleaning fluid, such as hydrogen water, principally for the removal of fine particles.

[0004] The inventor, et al. have found that the concurrent use of ultrasonic waves in the cleaning of bases by a reducing cleaning fluid resulted in extremely high levels of fine particle removal. However, this method, used on bare silicon surface that has been stripped off an oxidation film, has a disadvantage in that the base surface roughness increases when the strength of ultrasonic waves or the cleaning time is improper, such as when cleaning is performed over a lengthy period of time with a reducing cleaning water to which large ultrasonic waves are applied. In particular, the tendency for an increase in roughness becomes pronounced when an alkaline reducing cleaning water is used.

[0005] In the actual cleaning of silicon base surface, holding down the roughness of the silicon surface or that of the silicon/silicon oxidation film interface on the order of atoms is as important as the complete removal of impurities.

[0006] This invention aims to offer a simple and easy cleaning method that can both remove metal, organic and fine particle contaminants on the surface of electronic components, in particular of silicon bases, and also control an increase in the roughness of the base surface on the order of atoms in the cleaning process.

[0007] The inventor, et al. discovered that there was little likelihood of increased surface roughness caused by ultrasonic cleaning with a reducing cleaning fluid to which a chemical solution has been added, as long as the silicon base surface is oxidized prior to cleaning for the purpose of preventing an increase in the roughness of the base surface and a drop in the quality of silicon bases resulting from ultrasonic cleaning that is either too strong or too long in said ultrasonic cleaning with a reducing cleaning fluid. There is no damage to the interface between silicon and silicon oxidation film, which plays an important role in the silicon base functions. Building on these findings, this invention was completed.

[0008] Specifically, this invention provides the methods described in the following paragraphs.

(1) A method of cleaning electronic components characterized by cleaning of the components with an oxidizing cleaning fluid, followed by cleaning with a reducing cleaning fluid in the presence of ultrasonic vibrations.

(2) The method of cleaning electronic components according to Paragraph (1), wherein said electronic components are bare silicon bases for semiconductors.

(3) The method of cleaning electronic components according to Paragraph (1), wherein said oxidizing cleaning fluid is either ozone water or ozone water to which either an acidic or alkaline chemical has been added.

(4) The method of cleaning electronic components according to Paragraph (1), wherein said oxidizing cleaning fluid is either oxygen water or oxygen water to which either an acidic or alkaline chemical has been added.

(5) The method of cleaning electronic components according to Paragraph (1), wherein said oxidizing cleaning fluid is a fluid which contains hydrogen peroxide

(6) The method of cleaning electronic components according to Paragraph (1), wherein said reducing cleaning fluid is either hydrogen water or hydrogen water to which either an acidic or alkaline chemical has been added.

(7) The method of cleaning electronic components according to Paragraph (1), wherein ultrasonic vibrations have frequencies ranging between 400 KHz and 3 MHz.

(8) The method of cleaning electronic components according to paragraph(3), wherein said ozone concentration in a cleaning fluid is not less than 0.1mg/liter.

(9) The method of cleaning electronic components according to paragraph(5), wherein said hydrogen peroxide concentration in a cleaning fluid is not less than 200 mg/liter.

(10) The method of cleaning electronic components according to paragraph(6), wherein said hydrogen gas concentration in a cleaning fluid is not less than 0.7mg/liter.

[0009] According to one aspect of this invention there is provided a method of cleaning electronic components with a reducing cleaning fluid in the presence of ultrasonic vibrations, following their cleaning with an oxidizing cleaning fluid, and involves a cleaning process in which components are cleaned using a reducing cleaning fluid in the presence

of ultrasonic vibrations, either directly following a cleaning process that employs an oxidizing cleaning fluid or via an ultrapure water rinsing process.

[0010] The cleaning method is especially applicable to electronic components which demand strict surface roughness precision, such as silicon bases for semiconductors, and glass bases for liquid crystals.

[0011] In particular, this invention allows silicon bases for semiconductors to have either a hydrophilic surface finish with an oxidized film on the surface, or a hydrophobic surface finish without an oxidized film. The cleaning method is especially suitable for bare silicon bases for semiconductors and restrains the occurrence of base surface roughness.

[0012] In an embodiment of this invention, the initial treatment with an oxidizing cleaning fluid oxidizes the silicon base surface prior to the cleaning process in which a reducing cleaning fluid is used in the presence of ultrasonic vibrations. For this reason, the silicon/silicon oxidation film interface does not get damaged in the second cleaning process. Thus, there is no increase in electronic component (silicon base) surface roughness following the cleaning treatment.

[0013] The oxidizing cleaning fluid may be an aqueous solution in which an oxidizing substance has been dissolved.

[0014] There is no special restriction on the oxidizing substance used in the oxidizing cleaning fluid. For example, oxidizing substances, such as hydrogen peroxide (H_2O_2), ozone (O_3) and oxygen (O_2); hypochlorites, such as sodium hypochlorite ($NaClO$) and calcium hypochlorite; chlorite, such as sodium chlorite ($NaClO_2$) and potassium chlorite; and chlorates, such as sodium chlorate ($NaClO_3$) and ammonium chlorate, can be mentioned. These oxidizing substances can be used either singularly or in combination of two or more.

[0015] Among these, hydrogen peroxide and ozone are particularly suitable because they are relatively easy to handle, achieve high cleaning effects at low concentration, and place only minor load on the rinse after cleaning. When hydrogen peroxide is used, it is desirable for the hydrogen peroxide concentration in the cleaning water to be not lower than 200 mg/liter, and more preferably not lower than 1,000 mg/liter. When ozone is used, the ozone concentration in the cleaning water should be not lower than 0.1 mg/liter, and more preferably not less than 1 mg/liter.

[0016] Furthermore, fluids which contain these oxidizing substances, supplemented with an acidic or alkaline chemical, can be used. In particular, fluids containing ozone water to which either an acidic or alkaline chemical is added, and fluids containing oxygen water to which an acidic or alkaline chemical is added are most suited.

[0017] The reducing cleaning fluid used in an embodiment of this invention is water in which a reducing substance is dissolved.

[0018] There is no special restriction on the reducing substance used here. For example, reducing gases, including hydrogen gas; hyposulfites, such as sodium hyposulfite ($Na_2S_2O_4$) and ammonium hyposulfite; sulfites, such as sodium sulfite (Na_2SO_3) and ammonium sulfite; hydrogensulfites, such as sodium hydrogensulfite ($NaHSO_3$), and ammonium hydrogensulfite; nitrites, such as sodium nitrite ($NaNO_2$) and ammonium nitrite; hydrazine; inorganic reducing substances, such as hydrogen sulfide; organic reducing substances, such as formic acid and aldehyde, can be mentioned. These reducing substances can be used either singularly or in combination of two or more. Of these, hydrogen water, in which hydrogen gas has been dissolved, is especially suitable because it achieves high cleaning effects at low concentration, and places only minor load on the rinse after cleaning. It is desirable for the concentration of hydrogen gas in the water used for cleaning electronic components to be not lower than 0.7 mg/liter, and more preferably not lower than 1 mg/liter.

[0019] An acidic or alkaline chemical may be added to the aqueous solution in which the said reducing substance is dissolved, as the reducing cleaning fluid.

[0020] This method can be highly effective in cleaning with hydrogen water to which an alkaline chemical that is effective in removing fine particles is added and used concurrently with ultrasonic waves.

[0021] The purity of water in which reducing substances or oxidizing substances are dissolved can be set to meet the surface cleanliness demanded of the item to be cleaned. In other words, cleaning water for each cleaning process is prepared by dissolving either a reducing substance or substances, or an oxidizing substance or substances in the water having a level of purity which is non-contaminated in relation to the level of the surface cleanliness of the item to be cleaned. The water is used in the process in which contaminants on the surface of the item are removed by allowing said water to come into contact with the item to be cleaned.

[0022] Consequently, in situations where the item to be cleaned is a simple component that does not require especially strict levels of cleanliness, tap water in which industrial-purpose reducing substances and oxidizing substances have been dissolved can be used as the water for cleaning electronic components.

[0023] However, when cleaning the surface of electronic components, such as silicon bases for semiconductors, glass bases for liquid crystals, quartz bases for photomasks and other precision electronic components, it is desirable to use ultrapure water having sufficiently high purity and dissolve reducing substances of high purity or oxidizing substances of high purity. It is desirable for ultrapure water to have an electric resistance rate not lower than 18 $M\Omega \cdot cm$ at 25°C, and contain not more than 10 $\mu g/liter$ of organic carbons, and not more than 10,000 particles per liter. Furthermore, minute foreign particles in the cleaning water for electronic components can be removed through a filter if necessary.

[0024] In the process of cleaning with a reducing cleaning fluid, ultrasonic waves are applied to the reducing cleaning fluid in order to facilitate cleaning. Ultrasonic vibrations are excellent as a supplementary cleaning tool because they do not damage the surface of electronic components.

[0025] There is no particular restriction as to the methods of applying ultrasonic waves to reducing cleaning fluids of this invention. For example, in batch cleaning, ultrasonic vibrations can be delivered to a tank in which cleaning fluid for electronic components is stored. In spin cleaning, ultrasonic vibrations can be delivered to the nozzle from which the cleaning fluid for electronic components is sprayed.

[0026] The frequency of the ultrasonic waves applied to the ultrasonic wave cleaning process is desired to be between 20 KHz and 3 MHz, and more preferably between 400 KHz and 3 MHz. Ultrasonic wave frequencies of below 20 KHz can result in inadequate removal of fine particles from electronic components which are contaminated with fine particles. Frequencies in excess of 3 MHz do not produce improved effects commensurate with the higher frequencies.

[0027] As the result of the treatment with oxidizing cleaning fluids described above, organic and metal contaminants can be removed from the surface of electronic components, and an oxidization film is formed on the surface of the components. In the ensuing cleaning with reducing cleaning water, fine particles which adhere to the surface of electronic components are removed. As the result of this series of cleaning processes employing oxidizing cleaning water and reducing cleaning water, all contaminants, including organic matter, metals and fine particles, are removed.

[0028] In the removal of fine particles that adhere to bare silicon, no increase occurs between the surface roughness of bare silicon, which is the raw material, and that of silicon bases after cleaning if components are pre-treated with an oxidizing fluid, followed by a reducing cleaning fluid unlike when the conventional cleaning method is used wherein components are cleaned directly with a reducing cleaning fluid.

[0029] Therefore, when there is a cleaning process which precedes the preparation of bare silicon bases from silicon which has an oxidization film, by means of a cleaning treatment with a hydrofluoric acid chemical, such as a dilute hydrofluoric acid (DHF) cleaning fluid; cleaning with the buffered hydrofluoric acid (dilute hydrofluoric acid and ammonium fluoride); or dry treatment with anhydrous hydrofluoric acid gas, it is desirable that cleaning of this invention is performed following the first cleaning.

[0030] If it is necessary to give a hydrophobic surface finish stripped off an oxidization film to silicon that has been cleaned by the cleaning method, cleaning with a chemical solution, such as that of DHF, that is capable of dissolving a silicon oxidization film will result in a silicon surface that retains low roughness as measured on the same atomic orders as the raw material silicon.

[0031] If, on the other hand, a hydrophilic surface finish with an oxidization film is desired, the silicon treated with the cleaning method can be used without further treatment. Furthermore, a low-roughness silicon oxidization film surface that reflects the roughness of the raw material silicon surface can also be obtained by applying an oxidization treatment with ozone water, etc. after an oxidization film is once dissolved with DHF as described above.

[Examples]

[0032] In the following section, this invention is explained in further detail by using examples. This invention, however, is not limited in any way by these examples.

[0033] In blank tests, examples and comparative examples, cleaning effects were evaluated using the following method:

(1) Item to be Cleaned

[0034] Bare silicon bases having a diameter of 6 in. were submerged in ultrapure water containing alumina particles having a diameter of not more than 1 μm and divalent copper ions for 3 minutes, rinsed with ultrapure water to create contaminated silicon bases. These prepared silicon bases were used as the items to be cleaned.

[0035] The contaminated silicon bases had between 20,000 and 25,000 particles per base, and adhered copper in the amounts of $1\text{E} + 14$ atoms/cm².

(2) Cleaning Operation

[0036] After the contaminated silicon bases were cleaned with 0.5% dilute hydrofluoric acid, they were cleaned with ozone water having the concentration of 5 ppm as an oxidizing cleaning fluid, followed by cleaning with hydrogen water having the concentration of 1.2 ppm under the application of ultrasonic vibrations by a nozzle that generates ultrasonic vibrations having the frequency of 1.6 MHz.

[0037] All cleaning operations were performed by spin cleaning at the rotation speed of 500 rpm.

[0038] Cleaning water used in the tests was supplied at 0.8 liter/min.

[0039] Treatment duration in individual processes in the examples and comparative examples were as follows:

Dilute hydrofluoric acid (0.5%) treatment process	1 minute
Ozone water (5 ppm) treatment process	1 minute
Hydrogen water (1.2 ppm) treatment process	0.2 or 2 minutes

[0040] The time for the hydrogen water treatment process was set longer in this test than the time required in normal cleaning in order to allow surface damage effects to emerge more readily.

[0041] Following each step and also after the final step, ultrapure water rinsing was performed for 10 seconds.

[0042] The following blank tests were conducted to provide criterion for test evaluation:

Blank Test No. 1	Ozone water cleaning only.
Blank Test No. 2	Ozone water cleaning + dilute hydrofluoric acid treatment
Blank Test No. 3	Hydrogen water cleaning (0.2 minute) - (A)
	Ammonia-added hydrogen water cleaning (0.2 minute) - (B)
Blank Test No. 4	Hydrogen water cleaning (2 minutes) - (A)
	Ammonia-added hydrogen water cleaning (2 minutes) - (B)

(3) Evaluation

(3-1) Fine Particle Removal Rate

[0043] Fine particles were measured using base surface foreign particle testing equipment by the laser scatter method. Particle removal rates were computed by comparing particle counts before and after cleaning.

(3-2) Copper Removal Rate

[0044] Concentration of copper on the surface was measured by the total reflection fluorescent X-ray spectroscopy. Copper removal rates were computed based on copper concentration before and after cleaning.

(3-3) Surface Roughness (Increases in irregularity at the atomic level)

[0045] Samples were cut to 10 mm squares. The maximum height/depth distance in the $5 \mu \times 5 \mu$ square at the center of the sample was measured by AFM (atomic force microscope). Standard untreated bare silicon values ranged between 23 and 25 nm. Those whose post-cleaning values were not more than 27 nm were evaluated to have had "no increase in the roughness". Those whose values exceeded 27 nm were evaluated to have had "an increase in the roughness".

Example 1

[0046] The above-mentioned bare silicon bases, which are the items to be cleaned, were cleaned with hydrogen water (1-2 ppm)(A) for 0.2 minutes, and hydrogen water (1.2 ppm) to which 1 ppm of ammonia had been added (B) for 0.2 minutes, under the application of ultrasonic waves, after they were treated in the above-described 1-minute dilute hydrofluoric acid treatment process and 1-minute ozone water treatment process.

[0047] The cleaned items were evaluated using the above-described evaluation method.

Example 2

[0048] Bases were cleaned under conditions identical to those in Example 1 with the exception of the hydrogen water treatment time, which was changed to 2 minutes.

Example 3

[0049] Silicon bases, cleaned by the same way as under Example 2, were treated with dilute hydrofluoric acid to give a hydrophobic surface finish, and evaluated.

Example 4

[0050] Silicon bases, cleaned and treated with dilute hydrofluoric acid to produce a hydrophobic surface finish as under Example 3, were treated with ozone water to create a hydrophilic surface finish, and evaluated.

Comparative Example 1

[0051] Altering the order of the cleaning processes in Example 1, a 1-minute dilute hydrofluoric acid treatment was followed first by a 0.2-minute hydrogen water treatment, and then a 1-minute ozone water treatment.

[0052] The conditions of individual treatment processes were held the same as in Example 1

Comparative Example 2

[0053] Altering the order of the cleaning processes in Example 2, a 1-minute dilute hydrofluoric acid treatment was followed first by a 2-minute hydrogen water treatment, and then a 1-minute ozone water treatment.

[0054] The conditions of individual treatment processes were held the same as in Example 2.

Comparative Example 3

[0055] Altering the order of the cleaning processes in Example 3, a 1-minute dilute hydrofluoric acid treatment was followed first by a 2-minute hydrogen water treatment, and then a 1-minute ozone water treatment. Additionally, a dilute hydrofluoric acid treatment was given.

[0056] The conditions of individual treatment processes were held the same as in Example 3.

Comparative Example 4

[0057] Altering the order of the cleaning processes in Example 4, a 1-minute dilute hydrofluoric acid treatment was followed first by a 2-minute hydrogen water treatment, and then a 1-minute ozone water treatment. Additionally, a dilute hydrofluoric acid treatment was given, followed by an ozone water treatment.

[0058] The conditions of individual treatment processes were held the same as in Example 4.

[0059] Table 1 shows a summary of the evaluation performed on the cleaning results of the silicon bases in the above-described blank tests, examples and comparative examples under their respective treatment conditions.

[Table 1]

	Surface Finish	Fine Particle Removal Rate (%)	Copper Removal Rate (%)	Increase in Roughness at the Atomic Level
Blank Test 1	Hydrophilic	10	99<	None
Blank Test 2	Hydrophobic	50	99<	None
Blank Test 3	A Hydrophilic	85	<10	None
	B Hydrophilic	96	<10	None
Blank Test 4	A Hydrophilic	99	<10	None
	B Hydrophilic	100	<10	None
Example 1	A Hydrophilic	89	99<	None
	B Hydrophilic	97	99<	None
Example 2	A Hydrophilic	99	99<	None
	B Hydrophilic	100	99<	None
Example 3	A Hydrophobic	99	99<	None
	B Hydrophobic	100	99<	None
Example 4	A Hydrophilic	99	99<	None
	B Hydrophilic	100	99<	None
Comparative Example 1	A Hydrophilic	84	99<	None
	B Hydrophilic	93	99<	None
Comparative Example 2	A Hydrophilic	99	99<	Increase (32 nm)
	B Hydrophilic	99	99<	Increase (43 nm)
Comparative Example 3	A Hydrophobic	99	99<	Increase (33 nm)
	B Hydrophobic	99	99<	Increase (44 nm)
Comparative Example 4	A Hydrophilic	99	99<	Increase (32 nm)
	B Hydrophilic	100	99<	Increase (41 nm)

[0060] Based on the results of this table, oxidization treatment given prior to ultrasonic cleaning by hydrogen water

ensures the prevention of an increase in roughness on the base surface on the order of atoms.

[0061] The cleaning method herein described not only achieves high cleaning effects with respect to metals, organic matter and fine particles but also enables the maintenance of roughness at low levels on the order of atoms.

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Claims

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1. A method of cleaning electronic components characterized by cleaning with a reducing cleaning fluid while the components are subjected to ultrasonic vibrations, after cleaning with an oxidizing cleaning fluid.

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2. The method of cleaning electronic components according to Claim 1, wherein said electronic components are bare silicon bases for semiconductors.

3. The method of cleaning electronic components according to Claim 1, wherein said oxidizing cleaning fluid is either ozone water or ozone water to which an acidic or alkaline chemical has been added.

4. The method of cleaning electronic components according to Claim 1, wherein said oxidizing cleaning fluid is either oxygen water or oxygen water to which an acidic or alkaline chemical has been added.

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5. The method of cleaning electronic components according to Claim 1, wherein said oxidizing cleaning fluid is a solution containing hydrogen peroxide.

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6. The method of cleaning electronic components according to Claim 1, wherein said reducing cleaning fluid is either hydrogen water or hydrogen water to which an acidic or alkaline chemical is added.

7. The method of cleaning electronic components according to Claim 1, wherein said ultrasonic vibrations have frequencies between 400 KHz and 3 MHz.

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8. The method of cleaning electronic components according to Claim 3, wherein said ozone concentration in a cleaning fluid is not less than 0.1 mg/liter.

9. The method of cleaning electronic components according to Claim 5, wherein said hydrogen peroxide concentration in a cleaning fluid is not less than 200 mg/liter.

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10. The method of cleaning electronic components according to Claim 6, wherein said hydrogen gas concentration in a cleaning fluid is not less than 0.7 mg/liter.

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